# **SKY/FISH of Previously G-Banded Slides**

## Section of Cancer Genomics, Genetics Branch, NCI National Institutes of Health

### **Reagents**

Acetic Acid, glacial
Deionized Formamide
Ambion, Cat. # 9342
dH<sub>2</sub>O
Ethanol, absolute
MgCl<sub>2</sub>, 1M
Methanol, anhydrous
Mallinckrodt AR (ACS), Cat. 3016
Phosphate Buffered Saline (PBS) 1X
Rubber Cement
SSC, 2X
Xylene

# Preparation

**Fixative (3:1** Methanol:Acetic Acid [Vol:Vol])

Methanol 45 ml Acetic Acid 15 ml

1X PBS/MgCl<sub>2</sub> (f.c. of MgCl<sub>2</sub> = 50mM)

1M MgCl<sub>2</sub> 50 ml 1X PBS 950 ml

#### 1% Formaldehyde /1XPBS/MgCl<sub>2</sub>

Formaldehyde (37%) 2.7 ml 1X PBS 97.3 ml

#### FA/SSC

Deionized Formamide (FA) 70 ml 2X SSC 30 ml

Adjust pH to 7.0

#### **Procedure: G-Band Slide Pretreatment**

- 1. If required for analysis, first image G-banded slides and record X and Y coordinates from the microscope so one can relocate the identical metaphase spreads that will be subsequently be hybridized with SKY probes. (see note 1).
- 2. Remove immersion oil, if any, from previously G-banded slide by washing the slide in a coplin jar containing xylene for 2 minutes.
- 3. Rinse slide in a 50 ml coplin jar containing methanol for 2 min.
- 4. De-stain slide by immersing into a 50 ml coplin jar containing fixative (Methanol:Acetic Acid).
- 5. Rinse slide in ddH<sub>2</sub>O twice for 5 min each
- 6. Rinse slide in 1X PBS twice for 5 min each
- 7. Wash slide in 1X PBS/MgCl<sub>2</sub> for 5 min
- Place slide in 50 ml coplin jar containing
   1% Formaldehyde/1X PBS/1M MgCl<sub>2</sub>, for 10 min.
- 9. Wash slide for 5 min in 1X PBS, shaking.
- 10. Dehydrate slide in ethanol series; 70%, 90%, 100% ethanol, 2 min each
- 11. Air dry slide

## **Procedure: G-Band Slide Denaturation and Hybridization**

- 1. First fill a 50 ml coplin jar with FA/SSC (pH 7, pre-warmed to 70 °C) which is then placed inside a waterbath set at 70 °C (see note 2).
- 2. To denature the slide, place it inside the coplin jar for approximately 10-30 sec. Quickly remove slide and place into a new coplin jar containing ice cold ethanol (70%) for 2 min, followed by two more washes in 90%, and 100% ethanol, for 2 min each; air dry slide.
- 3. Apply probe to dry slide, carefully place a glass coverslip (18 mm²) over the probe and seal all edges with rubber cement.

4. Hybridize slide for 48-72 hours at 37 °C in a moist light tight container (see note 3).

#### **Notes**

- 1. As these slides have been previously G-banded, the slide pre-treatment using pepsin is eliminated as the chromosomes have already been partially digested with trypsin. G-banding enhances the DAPI banding and often results in brighter hybridization signals than the slides pre-treated with pepsin.
- 2. The time of slide denaturation is dependent on the age of the slide and the extent of digestion by the enzyme (trypsin) used in the G-banding procedure. Older slides (>2 months) require longer denaturation times.
- 3. Procede with detection of SKY hybridization according to protocols outlined in <u>Detection for SKY located under SKY Protocols.</u>